Product Codes Making the Connection....

Julie "Brandi" Stuart
Center Product Code Coordinator
Consumer Safety Officer, 510(k) Staff
FDA Office of Evaluation, CDRH

Public Law 94-295 94th Congress

An Act

To amend the Federal Food, Drug, and Cosmetic Act to provide for the safety and effectiveness of medical devices intended for human use, and for other DUBINOSES.

May 28, 1976 [S. 510]

Be it enacted by the Senate and House of Representatives of the United States of America in Congress assembled,

Medical Device Amendments of 1976. 21 USC 301 note.

SHORT TITLE AND TABLE OF CONTENTS

SECTION 1. (a) This Act may be cited as the "Medical Device Amendments of 1976".

(b) Whenever in this Act (other than in section 3(a)(1)(B)) an amendment is expressed in terms of an amendment to a section or other provision, the reference sl.ll be cons or other provision of th

21 USC 301.

Device Amendments

(c) Classification panel organization and operation,

Classification. Classification changes Late 1 2 295

"Sec. 514. Performance standards.

"(a) Provisions of standards.

"(b) Initiation of a proceeding for a performance standard.

"(c) Invitation for standards.

"(d) Acceptance of certain existing standards.

"(e) Acceptance of offer to develop standard.

"(f) Development of standard by Secretary after publication of subsection (c) notice.

"(g) Establishment of a standard.

"Sec. 515. Premarket approval.

"(a) General requirement.

*(b) Regulation to require premarket approval.

"(c) Application for premarket approval.

"(d) Action on an application for premarket approval.

"(e) Withdrawal of approval of application.

"(f) Product development protocol.

"(g) Review.

"(h) Service of orders.

"Sec. 516. Banned devices.

"(a) General rule.

"(b) Special effective date.

"Sec. 517. Judicial review.

"(a) Application of section. "(b) Additional data, views, and arguments.

"(c) Standard for review. "(d) Finality of judgments.

Code of Federal Regulations (CFR)

The May 28, 1976, Medical Device Amendments required and led to: The classification of approximately 1,700 generic device types.

Code of Federal Regulations (CFR) 21 CFR Parts 862-892

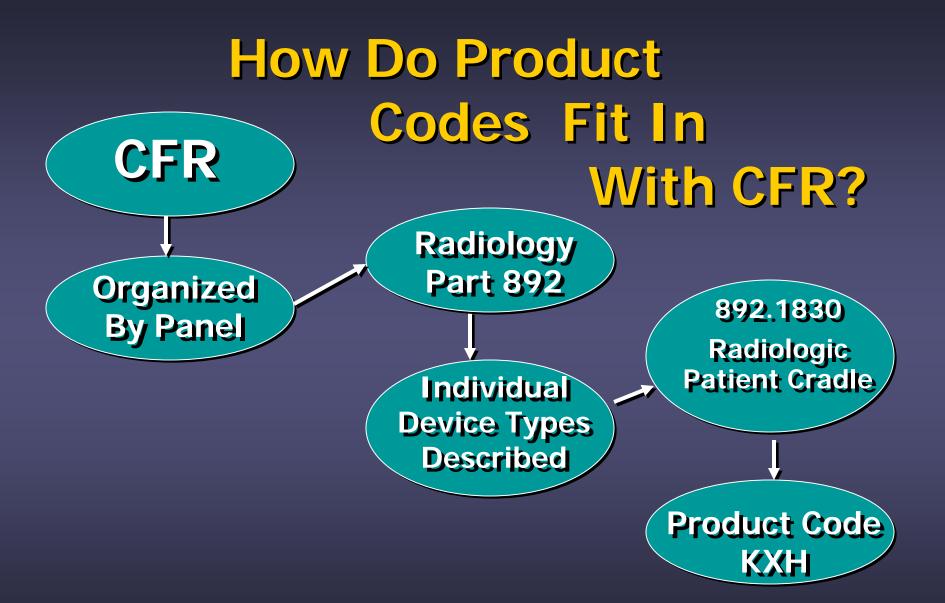


Code of Federal Regulations (CFR) 21 CFR Parts 862-892

Regulations describe the device types as they existed prior to May 28, 1976 (Pre-amendment)

May 28, 1976
Medical Device
Amendments
Public Law 94-295







U.S. Food and Drug Administration



CENTER FOR DEVICES AND RADIOLOGICAL HEALTH

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510(k) | Registration & Listing | Adverse Events | PMA | Classification | CLIA CFR Title 21 | Advisory Committees | Assembler | Recalls | Guidance | Standards

Radiology

Radiology

892.1830

510(k) Exempt

KXH

No

Plyluc Classification Database

Regular

Regulation Medical Specialty

Review Panel

Product Code

Submission Type

Regulation Number

Device Class GMP Exempt?

Note: FDA has exempted almost all class I devices (with the exception of Reserved Devices) from the premarket notification

Classification Today & Product Codes

Individual devices are classified by premarket review i.e., 510(k), PMA

New indications for use or new technologies are assigned new product codes that are placed under the original regulation

Preamendment Device Type Described in CFR

872.6865 — Powered Toothbrush

A powered toothbrush is an AC-powered or batterypowered device that consists of a handle containing a
motor that provides mechanical movement to a brush
intended to be applied to the teeth. The device is intended
to remove adherent plaque and food debris from the teeth
to reduce tooth decay.

Classification: Class 1 Exempt

Product Code Information: JEQ Powered Toothbrush Regulation: 876.6865 Class 1 Exempt

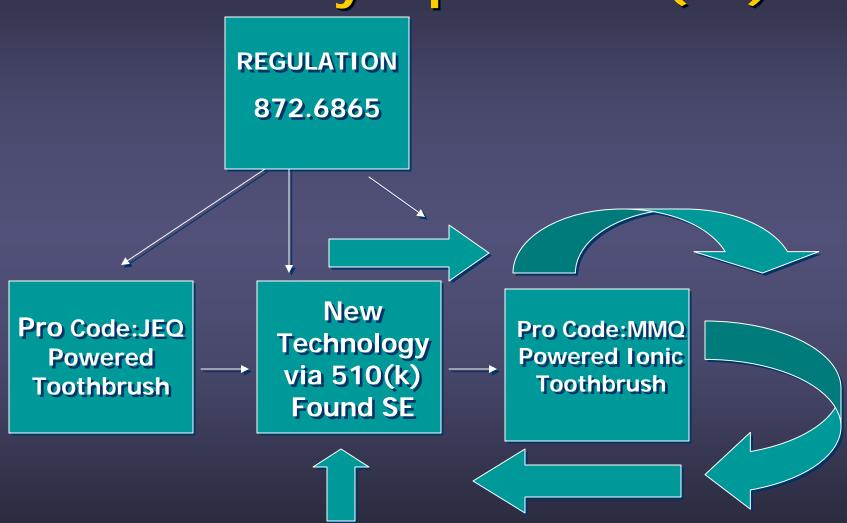
Substantially Equivalent

New Device Type: Ionic Powered Toothbrush
New Technology Required 510(k)
510(k) Found Substantially
Equivalent to Preamendment Device

Product Code Information: MMQ Powered Ionic toothbrush Regulation: 876.6865 Class 1 Exempt

▜

Substantially Equivalent (SE)



Why Are Product Codes Important To Me?

Ultimately classify the device Found on all 510(k) & PMA Letters

Tools:

- Required for Registration & Listing
- Used to Search for a Predicate
- Used to Search & Report Adverse Events
- Used Identify Third Party Eligible Device Types
- Required When Importing & Exporting Devices

Why Are Product Codes Important To Me?

Product Codes
Ultimately Classify
Your Device
Found on all 510(k)
and PMA Letters





Substantially Equivalent (SE)

Letter:

Food and Drug Administration 9200 Corporate Boulevard Rockville, Maryland 20850

Company ABC c/o John Doe 123 Street Name Somewhere, ST 99999

Product Codes are on all SE Letters and are available on the

Re: K078522

Internet Trade/Device Name: ABC Absorbable Gut Sutur

Regulation Number: 21 CFR 878.4830

Regulation Name: Absorbable surgical gut suture

Regulatory Class: II Product Code: GAK Dated: May 1, 2007 Received: May 2, 2007

Dear Mr. Doe:

Registration and Listing

A firm can ONLY list with the product code found on their 510(k) or PMA Clearance/Approval Letter

Unless registering with a Class 1 Exempt Device
Class 2 Exempt Device

Search For Predicates



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CFR Title 21 | Advisory Committees | Assembler | Recalls | Guidance | Standards

vww.acce	essdata.fda.gov/script
cdrh/cf	docs/cfPMM/pmn.cfm
Applicant Name	
Device Name	Expedited Review
Panel	Product Code
Decision	
Decision Date	to H
Sort by	Decision Date (descending)
	For full-text search, select Go To Simple Search button

Adverse Event Reporting



U.S. Food and Drug Administration



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- MAUDE data represents reports of adverse events involving medical devices. The data consists of voluntary reports since June 1993, user facility reports since 1991, di
 reports since 1993, and manufacturer reports since August 1996. MAUDE may not include reports made according to exemptions, variances, or alternative reporting requir
 granted under 21 CFR 803.19.
- The on-line search allows you to search CDRH database information on medical devices which may have malfunctioned or caused a death or serious injury. MAUDE is
 scheduled to be updated quarterly and the search page reflects the date of the most recent update. FDA seeks to include all reports received prior to the update. However,
 inclusion of some reports may be delayed by technical or clerical difficulties.
- MAUDE data is not intended to be used either to evaluate rates of adverse events or to compare adverse event occurrence rates across devices. Please be aware that regarding device vival vival vival across devices are rates across devices. Please be aware that regarding device vival viv

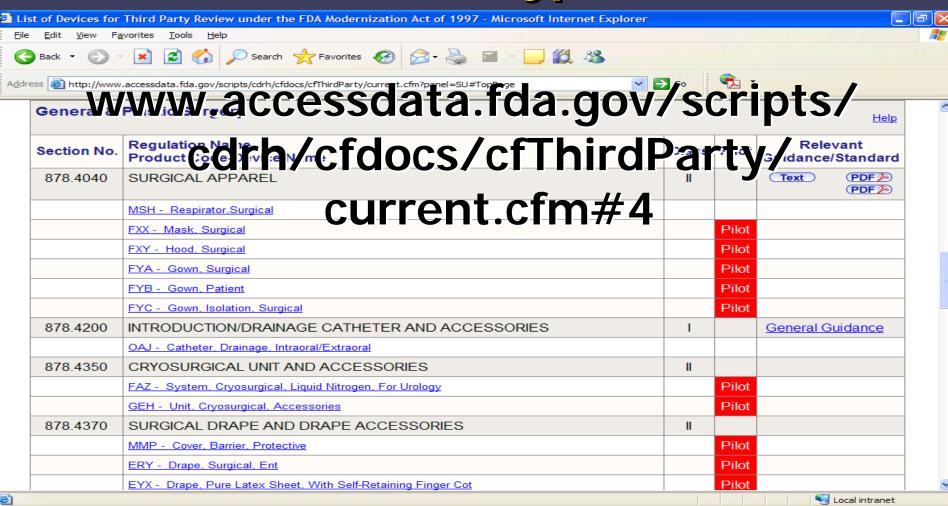
cdrh/efdocs/cfMAUDE/search.@FM...

Product Problem
Product Class
Brand Name
Manufacturer
Event Type

Date Report Received by FDA (mm/dd/yyyy)

For full-text search, select Go To Simple Search button

Search For Third Party Eligible Device Types



Microsoft O...

4 Internet...

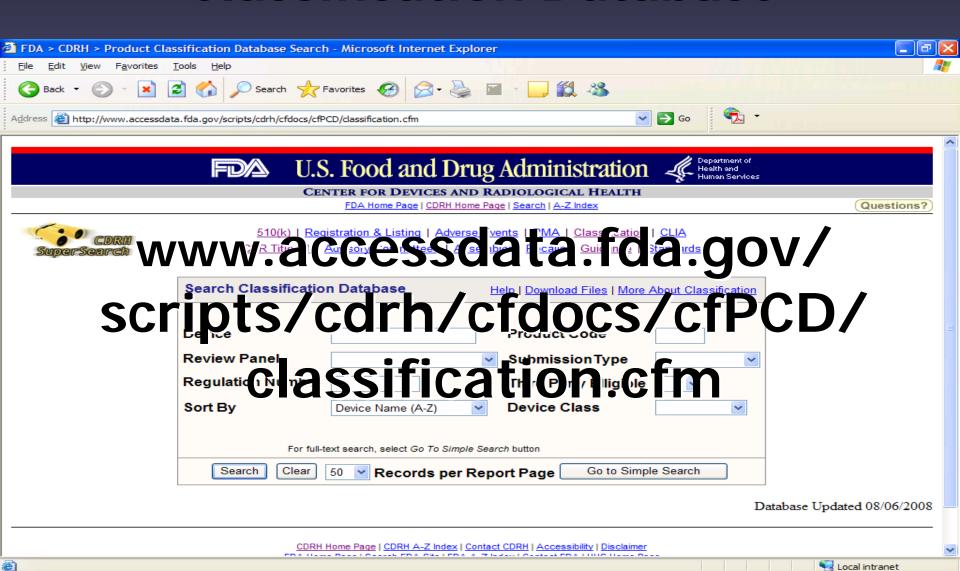
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Microsof...

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Classification Database



Product Code Descriptions

New Search

Back To Search Results

Product Classification Database

Device Elisa, Antibody, West Nile Virus

Regulation Description West Nile virus serological reagents.

Definition The west nile virus elisa is intended for the

detection of igg and igm antibodies to west nile virus. Specimens may be serum or cerebral spinal fluid from symptomatic

patients.

Regulation Medical

Specialty

Microbiology

Review Panel Microbiology

Product Code NOP

Submission Type 510(k)

Regulation Number 866.3940

Device Class 2

GMP Exempt? No

Guidance Document

Classification Database Classification Database

Provides:

Links to Standards;

Links to Associated Guidance Documents;

Product Code Definitions and Indications for Use Fields;

Expanded Descriptions; and

Third Party Eligibility

